e may be due to a system problem. Please contact your local STN Help Desk if you need assistance.

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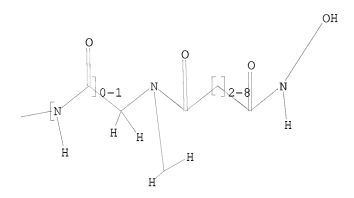
Uploading C:\Program Files\Stnexp\Queries\10580480c.str

STRUCTURE UPLOADED L8

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L8 HAS NO ANSWERS

Г8 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 18 sss

SAMPLE SEARCH INITIATED 14:35:36 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -2780 TO ITERATE

71.9% PROCESSED 2000 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

> **COMPLETE** BATCH

PROJECTED ITERATIONS: 52438 TO 58762 PROJECTED ANSWERS: 3 TO 205

L9 3 SEA SSS SAM L8

=> s 18 sss full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 191.05 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y FULL SEARCH INITIATED 14:35:50 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED -55174 TO ITERATE

100.0% PROCESSED 55174 ITERATIONS

82 ANSWERS

3 ANSWERS

SEARCH TIME: 00.00.02

08/09/2010 TOh

L10 82 SEA SSS FUL L8

=> file caplus
COST IN U.S. DOLLARS

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 194.97 410.54

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION

CA SUBSCRIBER PRICE 0.00 -2.55

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FILE COVERS 1907 - 8 Sep 2010 VOL 153 ISS 11

FILE LAST UPDATED: 7 Sep 2010 (20100907/ED)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Jun 2010

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2010

CAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2010.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 110

L11 18 L10

=> s 111 and Py2004 0 PY2004

L12 0 L11 AND PY2004

=> s 111 and Py<2004 24051605 PY<2004

L13 8 L11 AND PY<2004

=> s 111 and Py<2003

22999285 PY<20**0**3

L14 7 L11 AND PY<2003

=> d 113 1-8 ibib abs hitstr

THE ESTIMATED COST FOR THIS REQUEST IS 46.48 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:v

L13 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2003:485895 CAPLUS

DOCUMENT NUMBER: 139:223711

TITLE: Novel inhibitors of procollagen C-Proteinase. Part 2:

glutamic acid hydroxamates

AUTHOR(S): Robinson, L. A.; Wilson, D. M.; Delaet, N. G. J.;

Bradley, E. K.; Dankwardt, S. M.; Campbell, J. A.; Martin, R. L.; Van Wart, H. E.; Walker, K. A. M.;

Sullivan, R. W.

CORPORATE SOURCE: CombiChem Inc., San Diego, CA, 92121, USA

SOURCE: Bioorganic & Medicinal Chemistry Letters (2003

), 13(14), 2381-2384

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:223711

AB Glutamic acid derived hydroxamates were identified as potent and selective inhibitors of procollagen C-proteinase, an essential enzyme for the processing of procollagens to fibrillar collagens. Such compds. have

potential therapeutic application in the treatment of fibrosis.

IT 279255-52-6P 591766-04-0P 591766-06-2P

591766-07-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and structure-activity relationship of glutamic acid hydroxamates as novel inhibitors of procollagen C-Proteinase)

RN 279255-52-6 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)](4-methoxyphenyl)sulfonyl]amino]-N5-(2-cyanoethyl)-N1-hydroxy-N5-(phenylmethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 591766-04-0 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)[(4-

methoxyphenyl) sulfonyl]amino]-N5,N5-diethyl-N1-hydroxy-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 591766-06-2 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)](4-methoxyphenyl)sulfonyl]amino]-N1-hydroxy-N5-(2-phenylethyl)-N5-(phenylmethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 591766-07-3 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)](4-methoxyphenyl)sulfonyl]amino]-N5-(2-cyanoethyl)-N1-hydroxy-N5-(3-pyridinylmethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 13 THERE ARE 13 CAPLUS RECORDS THAT CITE THIS

RECORD (13 CITINGS)

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2002:275960 CAPLUS

DOCUMENT NUMBER: 136:310184

TITLE: Preparation of hydroxamic acid peptide deformylase

inhibitors as antibacterial agents

INVENTOR(S): Chong, Lee; Frechette, Roger; Scott, Carole; Tester,

Richard; Smith, Whitney; Chiba, Katsumi; Sakamoto,

Masatoshi; Gluchowski, Charles

PATENT ASSIGNEE(S): Questcor Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 171 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

TOh

PA	PATENT NO.				KIND DATE			APPLICATION NO.						DATE				
· · · -	WO 2002028829 WO 2002028829			A2 20020411 A3 20031224			WO 2001-US29926					20010924 <						
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DΖ,	EC,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	ΙL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	KΖ,	LC,	LK,	LR,	
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	NZ,	PH,	PL,	
		PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	
		UZ,	VN,	YU,	ZA,	ZW												
	RW:	GH,	GM,	ΚE,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AM,	ΑZ,	BY,	KG,	
		KΖ,	MD,	RU,	ΤJ,	TM,	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	
		ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	
		GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG									
AU	2002	0303	85		A 20020415				AU 2002-30385						20010924 <			
PRIORIT	Y APP	LN.	INFO	.:						US 2	000-	2349	67P		P 20000925			
										US 2	001-	7618	50		A 2	0010	118	
										WO 2	001-	US29	926	1	W 2	00109	924	
OTHER S GI	OURCE	(S):			MAR	PAT	136:	3101	84									

08/09/2010

AΒ Hydroxamic acid derivs. of peptides and peptidomimetics of formulas I, II, and III [wherein Z = NHOH or ORa; Ra = alkyl or a biocleavable moiety; X = CO or SO2; Y = (un)substituted heteroalkyl or heterocyclyl; R1 = (un) substituted (cyclo) alkyl, aryl, heterocyclyl, or heteroalkyl; R2R3 = 4-7 membered (un) substituted heterocycle; R2R4 = ring formed through a CH2CH2 linkage; or R2 = Me; or R3 = H or (un)substituted (hetero)alkyl, aryl, or heterocyclyl; or R4 = H or (un)substituted (hetero)alkyl, aryl, or heterocyclyl; R5 and R6 = independently H, NO2, NH2, NHCOH, NHCOCH3, NHSO2CH3, or (un) substituted CH2NH-(hetero)alkyl or CH2NH-heterocyclyl; one of R7 or R8 = CHR10CONHOH; one of R7 or R8 = (un)substituted (hetero)alkyl, (alkyl)heterocyclyl, or alkylaryl; R9 and R10 = $\frac{1}{2}$ independently H or (un) substituted (hetero) alkyl, (alkyl) heterocyclyl, or alkylaryl] were prepared as peptide deformylase (Fe-PDF) inhibitors for treating various bacterial infections. For example, 3-pyrrolidinol was added to tert-Bu (R)-(2-pentyl) succinate mono(N-hydroxysuccinimide) ester to give the amide (68%). Treatment with 20% TFA/DCM, followed by MeOH, benzene, and TMSN2 in hexanes, to afford the Me ester (90%). The pyrrolidinol was coupled with 4-methoxyphenylisocyanate and the ester converted to the hydroxamic acid (IV) using NH2OH⊕HCl. The latter inhibited E. coli Fe-PDF with IC50 of 9 nM and showed selectivity for Fe-PDF vs. thermolysin with a selectivity index of 30,000. Thus, I, II, and III are useful as antibiotics against a broad range of infectious disease in animals and humans.

IT 409129-80-2P 409129-81-3P 409129-82-4P

409129-83-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(peptide deformylase inhibitor; preparation of hydroxamic acid derivs. of peptides and peptidomimetics as peptide deformylase inhibitors for treatment of infectious diseases)

RN 409129-80-2 CAPLUS

CN Butanediamide, N4-hydroxy-N1-(2-hydroxyethyl)-2-pentyl-N1-(phenylmethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

Me
$$(CH_2)_4$$
 R N OH N OH

RN 409129-81-3 CAPLUS

CN Butanediamide, N4-hydroxy-N1, N1-bis(2-hydroxyethyl)-2-pentyl-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 409129-82-4 CAPLUS

CN Butanediamide, N1-[2-(3,4-dimethoxyphenyl)ethyl]-N4-hydroxy-N1-methyl-2-pentyl-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 409129-83-5 CAPLUS

CN Butanediamide, N4-hydroxy-N1-(2-hydroxyethyl)-N1-methyl-2-pentyl-, (2R)-(CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 11 THERE ARE 11 CAPLUS RECORDS THAT CITE THIS

RECORD (11 CITINGS)

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2001:453016 CAPLUS

DOCUMENT NUMBER: 135:61071

TITLE: Preparation of hydroxamic acid derivatives as matrix

metalloproteinase (MMP) inhibitors

INVENTOR(S): Owen, David Alan; Baxter, Andrew Douglas; Watson,

Robert John; Montana, John Gary

PATENT ASSIGNEE(S): Darwin Discovery Ltd., UK

SOURCE: PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA'	PATENT NO.			KIND DATE				APPLICATION NO.						DATE			
WO	2001	0441	 88		A1	_	2001	0621		wo 2	000-	GB48	61		2	0001	218 <
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		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	ΕE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
		HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KΡ,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
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		DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
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AU	2001	0220	17		A		2001	0625		AU 2	001-	2201	7		2	0001	218 <
EP	1237	8 6 7			A1		2002	0911		EP 2	000-	9856	09		2	0001	218 <
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL,	TR						
US	6462	042			В1		2002	1008		US 2	001-	8062	66		2	0010	328 <
PRIORIT	Y APP	LN.	INFO	.:					1	GB 19	999-	2997	9		A 1	9991	217
									,	WO 2	000-	GB48	61	,	W 2	0001	218
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OTHER SOURCE(S): MARPAT 135:61071

AB The title compds. B1NB2COCH2CR1R2CONHOH [I; R1 = alkyl, alkenyl, aryl, etc.; R2 = H, alkyl; CR1R2 = (un)substituted cycloalkyl, heterocycloalkyl; B1, B2 = H, alkyl, aryl, etc.] having therapeutic utility, were prepared E.g., a multi-step synthesis of (2S)-I [R1 = iso-Pr; R2 = H; B1 = Me; B2 = 4-(morpholin-4-yl)phenyl] was given. Compds. I are effective in treating inflammation at 0.01-50 mg/kg/day.

IT 345633-03-6P 345633-08-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of hydroxamic acid derivs. as matrix metalloproteinase (MMP) inhibitors)

RN 345633-03-6 CAPLUS

CN Butanediamide, N4-[2-(4-chlorophenoxy)ethy1]-N1-hydroxy-N4-methy1-2-(1-methylethy1)- (CA INDEX NAME)

RN 345633-08-1 CAPLUS

CN Butanediamide, N4-[3-(4-chlorophenyl)propyl]-N1-hydroxy-N4-methyl-2-(1-methylethyl)- (CA INDEX NAME)

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD

(7 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2000:441768 CAPLUS

DOCUMENT NUMBER: 133:74324

TITLE: Preparation of amino acid sulfonamide hydroxamates as

inhibitors of procollagen C-proteinase.

INVENTOR(S): Billedeau, Roland Joseph; Broka, Chris Allen;

Campbell, Jeffrey Allen; Chen, Jian Jeffrey; Dankwardt, Sharon Marie; Delaet, Nancy; Robinson,

Leslie Ann; Walker, Keith Adrian Murray

PATENT ASSIGNEE(S): F. Hoffmann-La Roche A.-G., Switz.

SOURCE: PCT Int. Appl., 133 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIND DATE			APPLICATION NO.						DATE					
WO.	20000374	36							 √() 1	aaa_	EDGG.	20		-	9991	214	/
NO	W: AE,												СН				
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	2355902			A1		2000									9991		
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AT	270271			Т		2004	0715	2	AT 1	999-	9635	30		1	9991	214	
RU	2232751			C2		2004				001-				1	9991	214	
US	6492394			В1		2002	1210	Ţ	JS 1	999-	4696	60		1	9991	222	<
	20010004	43		A2		2002				001-					20010		
	20010050			A		2002				001-					20010		
	20010063			A		2001				001-					20010		
	20010031			A		2001				001-					20010		
	20030199			A1		2003				002-					20021	-	
	6844366	J20		B2		2005		•	JU 2	002	2012.	<i>J</i>		2	.0021	005	
	20030216	105		A1		2003		1	TC 2	002-	2677	27		,	20021	000	/
	6787559			B2		2003	-	,	J	002-	20//	Z /		2	.0021	009	
	APPLN.			DZ		2004	0907		TC 1	998-	1122	11D		D 1	9981	222	
PRIORII.	APPLN.	INFO.	:												.9990		
															9991		
										999-					.9991		
										999-				-	9991	222	
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CHITED SO	MIRCE (S) .			MZDI	D A T	ા વવ∗	7/132	/1									

OTHER SOURCE(S): MARPAT 133:74324

HOHNCOCHRINRSO2Ar2 [R1 = alkyl, haloalkyl, heteroalkyl, cycloalkyl, aryl, aralkyl, aralkenyl, heteroaryl, heteroaralkyl, aminl, aryl, aralkyl, etc.; R = CHR2Ar1, CHR2CH: CHAr1; Ar2 = specified (substituted) Ph, naphthyl; R2 = H, alkyl; with provisos], were prepared Thus,

N-hydroxy-2(R)-[(3,4-methylenedioxybenzyl)(4-methoxy-2,3,6-

trimethylbenzenesulfonyl)amino]-3-methylbutyramide was prepared by solution

08/09/2010 TOh

phase synthesis from BOC-D-Val-OH. Title compds. inhibited procollagen C-proteinase with IC50 0.01-2 $\mu M.\,$

IT 279255-20-8P 279255-52-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amino acid sulfonamide hydroxamates as inhibitors of procollagen C-proteinase)

RN 279255-20-8 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)[(4-methoxy-2,3,6-trimethylphenyl)sulfonyl]amino]-N5-ethyl-N1-hydroxy-N5-methyl-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 279255-52-6 CAPLUS

CN Pentanediamide, 2-[(1,3-benzodioxol-5-ylmethyl)](4-methoxyphenyl)sulfonyl]amino]-N5-(2-cyanoethyl)-N1-hydroxy-N5-(phenylmethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS

RECORD (10 CITINGS)

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS

RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1998:498627 CAPLUS

DOCUMENT NUMBER: 129:175972

ORIGINAL REFERENCE NO.: 129:35769a,35772a

TITLE: Preparation of phenylsulfonamides as matrix

metalloproteinase inhibitors for treatment of diseases

INVENTOR(S): Takahashi, Kanji; Sugiura, Tsuneyuki

PATENT ASSIGNEE(S): Ono Pharmaceutical Co., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 42 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 10204054	A	19980804	JP 1997-20880	19970121 <
PRIORITY APPLN. INFO.:			JP 1997-20880	19970121
OTHER SOURCE(S):	MARPAT	129:175972		

GΙ

Phenylsulfonamides I [R1 = H, C1-4 alkyl; R2 = CO2R6, CONHOR7; R6, R7 = H, AΒ (un) substituted alkyl, Ph; R3 = OR11, (un) substituted amino, CO2R14, etc.; R11 = H, (un) substituted C1-4 alkyl, C2-4 acyl, etc; R14 = H, (un) substituted C1-4 alkyl, Ph; R4, R5 = H, (un) substituted C1-8 alky, (un) substituted amino, (hetero) cyclyl, etc.; E = CH:CH, C.tplbond.C; J = bond, C1-8 alkylene; R25 = H, (Ph-substituted) C1-4 alkyl, (Ph-substituted) alkoxycarbonyl] or their nontoxic salts are prepared The phenylsulfonamides are useful for treatment of rheumatoid arthritis, bone diseases, arteriosclerosis, tumor, autoimmune diseases, etc., caused by excess secretion or elevated activity of matrix metalloproteinase. Hydrolysis of N-[4-(4-hydroxy-1-butynyl)phenylsulfonyl]-D-tryptophan Me ester with aqueous NaOH gave 29% N-[4-(4-hydroxy-1-butyny1)phenylsulfonyl]-Dtryptophan, which inhibited gelatinase A activity at IC50 of 0.0079 μM. 211383-80-1P ΙT

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of phenylsulfonamides as matrix metalloproteinase inhibitors for treatment of diseases)

RN 211383-80-1 CAPLUS

CN Pentanediamide, N1-hydroxy-2-[[[4-(4-hydroxy-1-butyn-1-yl)phenyl]sulfonyl]amino]-N5-methyl-N5-(2-phenylethyl)-, (2R)- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L13 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1997:805715 CAPLUS

DOCUMENT NUMBER: 128:61793

ORIGINAL REFERENCE NO.: 128:12110h,12111a

TITLE: Preparation of N-(phenylsulfonyl)amino acid

derivatives as matrix metalloproteinase inhibitors

INVENTOR(S): Takahashi, Kanji; Sugiura, Tsuneyuki PATENT ASSIGNEE(S): Ono Pharmaceutical Co., Ltd., Japan

SOURCE: PCT Int. Appl., 194 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE			
WO 9745402			19970523 <			
RW: AT, BE, CH,		R, GB, GR, IE, IT,	LU, MC, NL, PT, SE			
JP 10265452	A 19981006	JP 1997-148448	19970523 < 19970523 <			
	A1 19990512 DE, DK, ES, FR, G		19970523 < NL, SE, PT, IE, FI			
PRIORITY APPLN. INFO.:		JP 1996-151864 JP 1997-20879	A 19970121			
OFFIED COUDON (C)	MADDAE 100 (170)	WO 1997-JP1735	W 19970523			

OTHER SOURCE(S): MARPAT 128:61793

GI

$$\begin{array}{c} \text{R}^1 \\ \text{SO}_2 \text{NR}^2 \text{OCR}^9 \text{R}^1 \text{O}_R \text{2} \\ \text{A-J-E} \end{array}$$

Me
$$C\equiv C$$
 SO_2-N CO_2R II

AΒ Phenylsulfonylamide derivs. represented by general formula (I; R1 = hydrogen or alkyl; R2 = CO2R3 or CONHOR4; wherein R3 = H, C1-8 alkyl, Ph, substituted C1-4 alkyl; R4 = H, C1-8 alkyl, Ph, phenyl-C1-4 alkyl; E = CH:CH, C.tplbond.C; A = hydrogen, alkyl, (un)substituted carbocycle or heterocycle; J = single bond or alkylene; R9, R10 = each hydrogen, (substituted) alkyl, COR11, carbocycle, heterocycle, etc.; R11 = OH, C1-8 alkyl, C1-8 alkoxy, PhO, phenyl-C1-4 alkyl, (un)substituted NH2; R20 = hydrogen, (substituted) C1-4 alkyl, C1-8 alkoxycarbonyl, phenyl-C1-4 alkoxycarbonyl, substituted C1-8 alkyl; or NR20CR9 = 5- to 7-membered heterocyclic ring containing 1 N atom) and salts thereof are prepared Also claimed are processes for producing the same; a matrix metalloproteinase inhibitor containing the same; and medicines containing the same and serving as preventives and/or remedies for rheumatism, osteoarthritis, pathol. bone resorption, osteoporosis, periodontosis, interstitial nephritis, arteriosclerosis, pulmonary emphysema, hepatocirrhosis, corneal injury, diseases due to cancer cell metastasis, infiltration and proliferation, autoimmune diseases (such as Crohn's disease and Sjogren's disease), diseases due to leukocyte emigration or infiltration, and neovascularization. Thus, 4-bromobenzenesulfonyl chloride was added to a solution of tert-Bu D-phenylalaninate in pyridine under ice-cooling and the resulting mixture was stirred at room temperature for 1 h to give tert-Bu N-(4-bromophenylsulfonyl)-D-phenylalaninate. A mixture of the latter compound, 10% Pd-C, Ph3P, CuI, MeCN, and Et3N was refluxed for 3 h to give tert-Bu D-phenylalaninate derivative (II; R = tert-butyl) which was stirred at room temperature for 1 h to give II (R = H). A tablet and an ampule formulation

containing II (R = H) were prepared

IT 200294-53-7P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of N-(phenylsulfonyl)amino acid derivs. as matrix metalloproteinase inhibitors for disease treatment)

RN 200294-53-7 CAPLUS

CN Pentanediamide, N1-hydroxy-N5-methyl-2-[[[4-[2-(4-methylphenyl)ethynyl]phenyl]sulfonyl]amino]-N5-(2-phenylethyl)-, (2R)-(CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 21 THERE ARE 21 CAPLUS RECORDS THAT CITE THIS

RECORD (24 CITINGS)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1994:700765 CAPLUS

DOCUMENT NUMBER: 121:300765

ORIGINAL REFERENCE NO.: 121:55057a,55060a

TITLE: Preparation of oxoheterocyclyl-substituted hydroxamic

acid derivatives as collagenase inhibitors

INVENTOR(S): Broadhurst, Michael John; Brown, Paul Anthony;

Johnson, William Henry; Lawton, Geoffrey

PATENT ASSIGNEE(S): F. Hoffmann-La Roche A.-G., Switz.

SOURCE: Eur. Pat. Appl., 27 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT NO.		KINI) I	DATE	API	PLICAT	ION NO.	 D.	ATE	
	574758 574758		A1 B1		19931222 19980909		1993-	108628	 1	993052	8 <
	R: AT,	BE, CH	DE,								
US	5318964		A		19940607	US	1993-	66832	1	993052	4 <
	9339816		А	-	19931216	AU	1993-	39816	1	993052	6 <
AU	659555		В2	-	19950518						
AT	170840		T	_	19980915	AT	1993-	108628	1	993052	8 <
ES	2121896		Т3		19981216	ES	1993-	108628	1	993052	8 <
ZA	9303957		A	_	19931213	ZA	1993-	3957	1	993060	4 <
RO	112613		В3	_	19971128	RO	1993-	777	1	993060	4 <
CZ	283373		В6	_	19980415	CZ	1993-	1081	1	993060	4 <
IL	105921		A	_	19980104	IL	1993-	105921	1	993060	7 <
CA	2098168		A1	_	19931212	CA	1993-	2098168	1	993061	0 <
NO	9302117		A		19931213	NO	1993-	2117	1	993061	>
CN	1083062		A	_	19940302	CN	1993-	107239	1	993061	>
CN	1035616		С	_	19970813						
JP	06065196		A	-	19940308	JP	1993-	165228	1	993061	0 <
JP	07076210		В	_	19950816						
FI	109535		В1	2	20020830	FI	1993-	2692	1	993061	1 <
US	5447929		А	_	19950905	US	1994-	214895	1	994031	7 <
PRIORITY	APPLN. I	NFO.:				GB	1992-	12421	A 1	992061	1
						GB	1993-	5720	A 1	993031	9

US 1993-66832 A3 19930524

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 121:300765

AB R1(CH2)nCH(CONHOH)CH(CONR2R3)CHR4CR5R6CH2R7 (R1 = N-attached oxoheterocyclyl; R2 = alkyl; R3 = alkyl or aryl; NR2R3 = heterocyclyl; R4-R7 = H or Me; n = 1-4) were prepared Thus, (2R)-[(1R,S)-tert-butoxycarbonyl-2-phthalimidoethyl]-4-methylvaleric acid was amidated by 1-benzyloxycarbamoyl-(3S)-hexahydropyridazinecarboxylic acid and the product converted in 3 steps to title compound (R,S)-I which had IC50 of 1.2 nM against collagenase in vitro.

Ι

RN 159135-28-1 CAPLUS

CN Hexanamide, 1-(1,3-dihydro-1,3-dioxo-2H-isoindol-2-yl)-N,N-diethyl-N'-hydroxy-5-methyl- (CA INDEX NAME)

RN 159135-30-5 CAPLUS

CN Hexanamide, 1-(1,3-dihydro-1,3-dioxo-2H-isoindol-2-yl)-N-ethyl-N'-hydroxy-N,5-dimethyl- (CA INDEX NAME)

THERE ARE 30 CAPLUS RECORDS THAT CITE THIS OS.CITING REF COUNT: 30 RECORD (38 CITINGS)

L13 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1984:531205 CAPLUS

DOCUMENT NUMBER: 101:131205

ORIGINAL REFERENCE NO.: 101:19977a,19980a

TITLE: Role of complex formation during polycondensation of

activated N-hydroxysuccinimide esters with diamines

Katsarava, R. D.; Kharadze, D. P.; Avalishvili, L. M.; AUTHOR(S):

Zaalishvili, M. M.

CORPORATE SOURCE: Inst. Fiziol., Tbilisi, USSR

Vysokomolekulyarnye Soedineniya, Seriya A (SOURCE:

1984), 26(7), 1537-43 CODEN: VYSAAF; ISSN: 0507-5475

DOCUMENT TYPE: Journal LANGUAGE: Russian

GΙ

AΒ During polycondensation of diamines with the title esters (I, Z =alkylene, arylene), the N-hydroxysuccinimide (II) [6066-82-6] byproduct formed complexes with the diamines. During polycondensation of weakly reactive I (Z = arylene) with aliphatic diamines at moderate temps., the complexation retarded polycondensation and prevented formation of high-mol.-weight polyamides. The polymerization rate increased sharply at

temperature; however, side reactions also intensified. During reaction of highly reactive I (Z = alkylene), complexation had little influence on the polymerization

ΙT 91990-28-2P

RL: PREP (Preparation)

(formation and properties of, polycondensation of diamines with hydroxysuccinimide diesters in relation to)

RN91990-28-2 CAPLUS

CN Butanediamide, N1, N1-diethyl-N4-hydroxy- (CA INDEX NAME)

$$\begin{array}{c|c} \mathsf{O} & \mathsf{O} & \mathsf{O} \\ || & || \\ \mathsf{HO-NH-C-CH}_2-\mathsf{CH}_2-\mathsf{C-NEt}_2 \end{array}$$